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# Practical-scale tests of cryogenic molecular sieve for separating low-concentration hydrogen isotopes from helium

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#### Abstract

Earlier bench-scale work at the Tritium Systems Test Assembly (TSTA) at Los Alamos National Laboratory examined a number of adsorbents for their suitability for separating low-concentration hydrogen (no tritium) from helium. One of the effective adsorbents was Linde 5A molecular sieve. Recently, experiments including tritium were conducted using practical-scale adsorbers. These tests used existing cryogenic molecular sieve beds (CMSBs) which each contain about 1.6 kg of Linde 5A molecular sieve. They are part of the TSTA integrated tritium processing system. Gas was fed to each CMSB at about 13 SLPM (standard liters per minute) with a nominal composition of 99% He, 0.98% H<sub>2</sub> and 0.02% HT. In all cases, for an extended period of time, the beds allowed no detectable (via Raman spectroscopy) hydrogen isotopes to escape in the bed effluent. Thereafter, the hydrogen isotopes appeared in the bed exit with a relatively sharp breakthrough curve. It is concluded that cryogenic molecular sieve adsorption is a practical and effective means of separating low-concentration hydrogen isotopes from a helium carrier.

# 1. Introduction

There are a number of cases in fusion fuel processing where low-concentration hydrogen isotopes need to be separated from helium. Usually the helium is a purge gas used to move hydrogen isotopes from one location to another. One of the most notable applications is associated with removing tritium from a solid ceramic breeder. For some designs which have been considered, helium with about 1% protium is purged through the ceramic. The protium exchanges with tritium which has been bred in the solid. The resulting gas composed of helium ( $\sim$ 99%), protium ( $\sim$ 1%) and tritium ( $\sim$ 0.01%) flows out of the blanket and, for further processing, requires separation of the hydrogen isotopes and the helium.

Earlier bench-scale work [1] at the Tritium Systems Test Assembly (TSTA) at Los Alamos National Laboratory examined a number of adsorbents for their suitability for separating low-concentration hydrogen (no tritium) from helium. One of the effective adsorbents was Linde 5A molecular sieve. Practical-scale, tritium-compatible beds packed with this adsorbent already exist at TSTA. They have now been used to separate low-concentration hydrogen isotopes (Q<sub>2</sub>) from He. The purpose of this paper is to report results from these experiments.

## 2. Ceramic breeding blanket processing

Fig. 1 shows a system which could be used for processing a ceramic lithium breeding blanket. Other

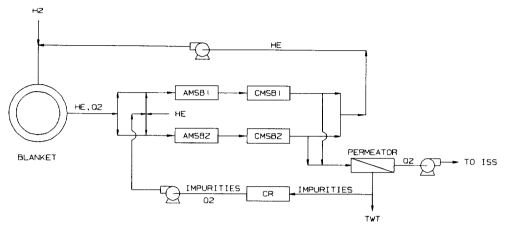


Fig. 1. Breeding blanket processing system proposed for ceramic lithium blanket processing. AMSB, ambient molecular sieve bed; CMSB, cryogenic molecular sieve bed; CR, catalytic reactor; ISS, isotope separation system; TWT, tritium waste treatment.

processes have also been considered [2,3]. For the process in Fig. 1, H<sub>2</sub> is added to He so that about 1% H<sub>2</sub> in He is swept through the blanket. The protium exchanges with tritium held in the blanket material so that the blanket exhaust is composed of He and a mixture of H<sub>2</sub> and HT. This stream may also contain some impurities, so the first processing step is an ambient temperature molecular sieve bed (AMSB), where impurities such as water are collected. Next is a cryogenic temperature molecular sieve bed (CMSB), where the remaining impurities and the hydrogen isotopes are collected. Leaving the CMSB is relatively pure He which is recycled back to the blanket. When the CMSB is saturated with Q<sub>2</sub> it is taken off-line for regeneration and its companion bed can be put into service. A CMSB is regenerated by warming. The Q<sub>2</sub> desorbs and is sent to a Pd/Ag permeator. The permeate from this device is ultrapure Q<sub>2</sub>, which can be sent directly to the isotope separation system (ISS). The retentate or "bleed" stream is sent to a shift catalyst bed where reactions such as steam reforming and the water gas shift reaction can be used to move hydrogen isotopes from impurities such as CQ4 and Q<sub>2</sub>O to the form of Q<sub>2</sub>. In place of the catalytic reactor an electrolysis system could be used to crack water directly if this were the only impurity present. The stream of Q<sub>2</sub> and impurities is recycled back through the regenerating CMSB train and over the permeator again. This circulation is continued until all of the Q2 has desorbed from the CMSB and all of the Q has been recovered from the impurities. The remaining tritiumfree impurities are exhausted to the tritium waste treatment (TWT) system.

# 3. Experimental

Within the various subsystems at TSTA exist almost all of the components necessary to demonstrate the process shown in Fig. 1. The Fuel CleanUp (FCU) system includes cryogenic molecular sieve beds (CMSBs) which each contain about 1.6 kg of Linde 5A molecular sieve. In the place of the AMSBs shown in Fig. 1, the FCU uses cold traps (freezers). A Pd permeator exists in the Japan Fuel CleanUp (JFCU) system. TSTA also has an isotope separation system consisting of four cryogenic distillation columns. All of the components are of ITER-relevant size. Tubing interconnects these components so they can be tested as an integrated system.

The actual set-up used for this experiment is shown in Fig. 2. Prior to the experiment, the CMSBs were regenerated at 500 K overnight with a helium purge. All lines and components were purged repeatedly with helium and evacuated to ensure that no residual impurities were present. The flows in lines 2 and 5 were controlled at 12.6 and 0.15 SLPM (standard liters per minute), respectively. The combination of these two flows were fed to one of the two CMSBs which were immersed in liquid nitrogen (77 K). Thus, the bed feed (line 1) flow rate was 12.75 SLPM with a nominal composition (before breakthrough) of 99% He, 0.98% H<sub>2</sub> and 0.02% HT. The gas leaving the CMSB (line 2) was recycled back to the CMSB feed using a metal bellows pump. A mixture of 2% HT in H<sub>2</sub> (line 5) was added to this recycle stream to maintain the desired bed feed composition. The gas composition at both the feed

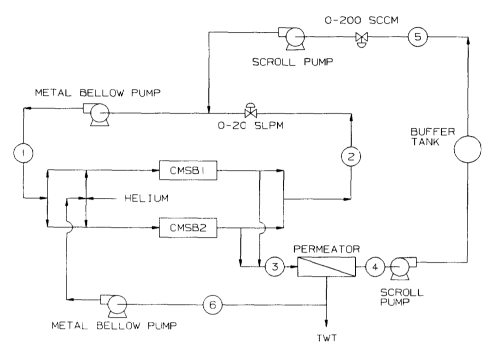


Fig. 2. Schematic diagram of run configuration.

and exit of the CMSBs was monitored alternately using on-line Raman spectroscopy.

After hydrogen isotopes appeared in the CMSB exit, the  $Q_2$  makeup (line 5) was turned off. Circulation was continued until a steady-state circulation loop composition was achieved. The equilibrium pressure and composition were recorded for comparison with previously determined isotherm values.

After equilibrium data had been recorded, the CMSB was warmed to about 500 K and the desorbed gas was sent to line 3 and was processed with the Pd diffuser. The ultrapure  $Q_2$  permeate (line 4) was stored in a buffer volume. The retentate or bleed (line 6) was recirculated back to the CMSB feed. Eventually almost all of the  $Q_2$  was removed from the circulation gas.

#### 4. Results and discussion

Three bed loading-regeneration cycles were completed during this campaign. Each cycle took one working day to complete.

For the first test cycle, CMSB1 was loaded. However, owing to time limitations, this bed loading was not done in a steady manner. Rather, the  $Q_2$  mixture was rapidly added to the line 1-2 circulation loop. The

results from this specific test were not meaningful for "breakthrough" (or mass transfer) information, and were only valuable for equilibrium information.

On the following two days, CMSB2 and CMSB1 were loaded in a controlled manner. The line 1-2 circulation loop was first filled with He and a flow rate of about 13 SLPM was maintained. To this loop, 0.15 SLPM of the  $Q_2$  mixture was added. Thus, the actual CMSB feed was about 1.15%  $Q_2$ . For both of these tests, gas analysis showed no detectable  $Q_2$  (about 100 ppm) at the CMSB exit for an extended period of time, i.e. more than 4 h. Thereafter, the  $Q_2$  at the bed exit was observed to rise rather rapidly and continued to rise until the  $Q_2$  makeup was stopped. Then the composition leveled off at a steady-state value.

Table 1 summarizes the equilibrium or isotherm information that can be drawn from these experiments. Using the equilibrium H<sub>2</sub> composition and total pressure, the H<sub>2</sub> partial pressure could be determined and the result recorded. Using pressure-volume-temperature measurements, the total amount of H<sub>2</sub> added during a run was calculated and listed. This value was divided by the approximate weight of sieve in each CMSB to determine the equilibrium bed loading and this result is given in Table 1.

Table 1 Summary of equilibrium results

No. Run	Bed	H <sub>2</sub> conen. (%)	HT conen. (%)	T <sub>2</sub> concn. (%)	Total pressure (Torr)	H <sub>2</sub> partial pressure (Torr)	H <sub>2</sub> adsorbed (std. l)	Bed loading (this work) (std. cm <sup>3</sup> H <sub>2</sub> g <sup>-1</sup> sieve)	Bed loading extrapolated from [1]
1	CMSB1	1.82	0.055	ND a	1159	21.1	56.3	35.2	51.7
2	CMSB1	1.68	0.030	ND	821	13.8	44.5	27.8	45.8
3	CMSB2	2.56	0.069	ND	672	17.2	45.7	28.6	49.0

<sup>&</sup>lt;sup>a</sup> ND means none detectable.

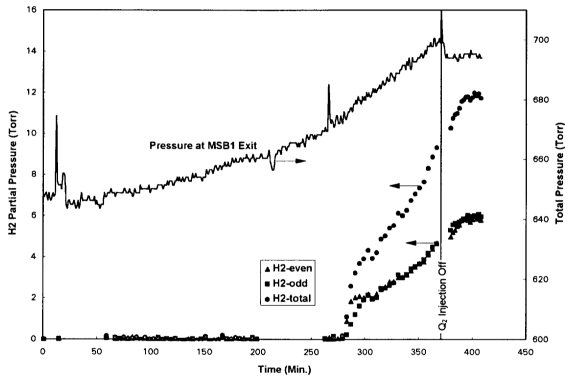


Fig. 3. Breakthrough curve for second CMSB1 loading (run 2).

For comparision purposes, the Langmuir isotherm equation determined for Linde 5A sieve previously [1] was used to calculate the bed loading that should be expected. This isotherm was shown to represent accurately the referenced bench-scale data. These values are much larger than those determined from the practical-scale beds. However, this is not entirely unexpected. The beds used in this work have been in place and in tritium service for about 8 years. There is uncertainty regarding the amount of sieve originally loaded into these beds and even more uncertainty after years of

service. These larger beds were not designed for high-temperature regeneration and residual material such as water on the beds would reduce their capacity. Hence it is not surprising that the fresh, well regenerated sieve used previously [1] would have a higher capacity. Owing to these uncertainties regarding the state of the practical-scale CMSBs, no definitive conclusions should be drawn from these data.

Breakthrough curves for the second CMSB1 loading (run 2) and the CMSB2 loading (run 3) are shown in Figs. 3 and 4, respectively. The total H<sub>2</sub> composition at

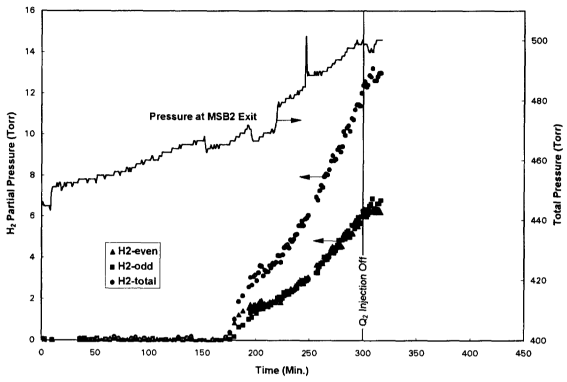


Fig. 4. Breakthrough curve for CMSB2 loading (run 3).

the bed exit and the bed exit pressure as a function of time after initiation of  $Q_2$  addition are shown. Also available from Raman analysis and shown in Figs. 3 and 4 are the fractions of  $H_2$  in the "odd" and "even" spin states. An examination of the first part of the breakthrough curves reveals that the "even"  $H_2$  breaks through first. Intermittently, analysis was also performed for HT and  $T_2$ , but the values were just at or below the Raman limits of detection (  $\sim 100$  ppm). Hence these data are not shown in Figs. 3 and 4.

As shown in Figs. 3 and 4, runs were conducted at different total pressures. This was by design so that the beds were exposed to different  $Q_2$  partial pressures. Prior to breakthrough, the pressure for run 2 increased from about 645 to 660 Torr (1 Torr = 133.3 Pa), while the corresponding increase for run 3 was from about 450 to 465 Torr. The feed composition was the same for both runs, so the partial pressure difference between the runs was simply proportional to the total pressure. At a higher partial pressure, beds should load with more  $Q_2$  and breakthrough should be delayed. Indeed, this was observed as the lower pressure

run 3 (Fig. 4) broke through after about 176 min, whereas the higher pres-sure run 2 (Fig. 3) broke through after about 280 min. At the initial point of breakthrough, the amounts of gas adsorbed were 42 and 26.4 standard liters for runs 2 and 3, respectively.

The single most remarkable result from these experiments is the relatively long time during which there is no detectable  $H_2$  at the bed exit, followed by a sharp  $Q_2$  breakthrough. A consideration in adsorber design is the "length of unused bed". This is a measure of the amount of unloaded adsorbent remaining when the adsorbate appears at the bed exit. It is apparent that for this work the length of unused bed is relatively short.

Further scrutiny of Figs. 3 and 4 discloses that the breaktrough curves have two distinct slopes. The first and more rapid rise has the shape that would be expected for a "once-through" experiment where the bed exhaust is not recycled back to the feed. However, this experiment does recycle the exhaust back to the feed. As breakthrough progresses, the recycled gas contains increasing amounts of  $Q_2$  that combines with the constant flow of  $Q_2$  makeup. The result is that the

 $Q_2$  composition or partial pressure in the bed feed increases. This, in turn, increases the equilibrium loading that the CMSB adsorbs. The second, less steep slope in Figs. 3 and 4 is indicative of this regime where the entire length of the bed is being loaded to higher adsorbed compositions.

Apparent in Figs. 3 and 4 are two distinct slopes for the total system pressure curves. The first, less steep, increase coincides with the period before  $H_2$  appears at the CMSB exit. This increase is interpreted to be due to additional helium filling the loop as it is being displaced from the CMSB by  $Q_2$ . After breakthrough, the total pressure increases more rapidly since the  $Q_2$  makeup is no longer all being adsorbed. When the  $Q_2$  injection is turned off, there is a brief equilibration period before the pressure reaches a steady-state value.

The pressures used to determine the equilibrium data reported in Table 1 are not shown in Figs. 3 and 4. Following the period of data collection shown on those figures, the circulation flow rate was decreased substantially so that pressure drops throughout the loop were minimal. Otherwise, a significant pressure drop would exist across the CMSB and the bed loading would vary along the length of the bed. The equilibrium pressures and compositions were measured after equilibration at the reduced flow rate.

During run 2, there is a gap in the  $H_2$  data between 200 and 260 min. At that time the Raman spectrometer was tuned to observe the HT concentration. There was none detectable. This indicates that HT did not break through before  $H_2$ . Indeed, this is the expected result based on chromatographic experiments which indicate that the order of elution for hydrogen isotopes is  $H_2$ , HD, HT,  $D_2$ , DT and  $T_2$ .

### 5. Conclusions

The most remarkable conclusion from this work is the relatively long period of time during which no Q2 was observed at the exit from the CMSB during its loading, followed by a sharp Q<sub>2</sub> breakthrough. This indicates that, even at low partial pressures (10-20 Torr), liquid nitrogen-cooled Linde 5A molecular sieve has considerable affinity and capacity for  $Q_2$ . It is therefore concluded that this material is appropriate for separating low-concentration Q<sub>2</sub> from He such as might be encountered when processing the effluent from a ceramic Li breeding blanket. This confirms that (except for the CR, which was not tested) the process shown in Fig. 1 is a workable solution for this application. There was nothing unexpected encountered in this work due to the presence of tritium compared with previous non-tritium experiments. No quantitative conclusions can be drawn from this work regarding equilibrium loading values for the adsorbent tested, but the numbers calculated are qualitatively in agreement with earlier experiments which were better suited for making these measurements.

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